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### Authors

Wagner, Sarah  
Roberson, David  
Boland, Joseph  
et al.

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# Evaluation of TypeSeq, a Novel High-Throughput, Low-Cost, Next-Generation Sequencing-Based Assay for Detection of 51 Human Papillomavirus Genotypes

Sarah Wagner,<sup>1,2</sup> David Roberson,<sup>1,2</sup> Joseph Boland,<sup>1,2</sup> Aimée R. Kreimer,<sup>1</sup> Meredith Yeager,<sup>1,2</sup> Michael Cullen,<sup>1,2</sup> Lisa Mirabello,<sup>1</sup> S. Terence Dunn,<sup>3</sup> Joan Walker,<sup>3</sup> Rosemary Zuna,<sup>3</sup> Carolina Porras,<sup>4</sup> Bernal Cortes,<sup>4</sup> Joshua Sampson,<sup>1,5</sup> Rolando Herrero,<sup>5</sup> Ana Cecilia Rodriguez,<sup>4</sup> Wim Quint,<sup>6</sup> Leen-Jan Van Doorn,<sup>6</sup> The CVT Group, Allan Hildesheim,<sup>1</sup> Mark Schiffman,<sup>1</sup> and Nicolas Wentzensen<sup>1,\*</sup>

<sup>1</sup>Division of Cancer Epidemiology and Genetics, National Cancer Institute, Bethesda, Maryland; <sup>2</sup>Cancer Genomics Research Laboratory, Frederick National Laboratory for Cancer Research, Leidos Biomedical Research Inc., Maryland; <sup>3</sup>University of Oklahoma Health Sciences Center, Oklahoma City; <sup>4</sup>Agencia Costarricense de Investigaciones Biomédicas, San José, Costa Rica; <sup>5</sup>International Agency for Research on Cancer, World Health Organization, Lyon, France; <sup>6</sup>DDL Diagnostic Laboratory, Rijswijk, The Netherlands

**Background.** Human papillomaviruses (HPV) cause over 500 000 cervical cancers each year, most of which occur in low-resource settings. Human papillomavirus genotyping is important to study natural history and vaccine efficacy. We evaluated TypeSeq, a novel, next-generation, sequencing-based assay that detects 51 HPV genotypes, in 2 large international epidemiologic studies.

**Methods.** TypeSeq was evaluated in 2804 cervical specimens from the Study to Understand Cervical Cancer Endpoints and Early Determinants (SUCCEED) and in 2357 specimens from the Costa Rica Vaccine Trial (CVT). Positive agreement and risks of precancer for individual genotypes were calculated for TypeSeq in comparison to Linear Array (SUCCEED). In CVT, positive agreement and vaccine efficacy were calculated for TypeSeq and SPF10-LiPA.

**Results.** We observed high overall and positive agreement for most genotypes between TypeSeq and Linear Array in SUCCEED and SPF10-LiPA in CVT. There was no significant difference in risk of precancer between TypeSeq and Linear Array in SUCCEED or in estimates of vaccine efficacy between TypeSeq and SPF10-LiPA in CVT.

**Conclusions.** The agreement of TypeSeq with Linear Array and SPF10-LiPA, 2 well established standards for HPV genotyping, demonstrates its high accuracy. TypeSeq provides high-throughput, affordable HPV genotyping for world-wide studies of cervical precancer risk and of HPV vaccine efficacy.

**Keywords.** HPV; TypeSeq; vaccine efficacy; genotyping; screening.

Human papillomaviruses (HPV) are a major cause of invasive cancers world-wide, with up to 600 000 cancers related to HPV occurring in 2012, 500 000 of which were cervical cancers. Less common HPV-related cancer sites include the vulva, vagina, anus, penis, and oral cavity [1]. Over 200 HPV genotypes have been identified so far that vary by tissue tropism and carcinogenic potential. Only a small group of 12–13 types is responsible for most HPV-related cancers (referred to as high

risk [HR] or carcinogenic types). Risk of precancer and cancer varies substantially within this group. HPV16 is by far the most carcinogenic type, causing over 60% of invasive cervical cancers and most HPV-related noncervical cancers, followed by HPV18 (15% of cervical cancers). At the cervix, both HPV16 and HPV18 increase in relative prevalence from HPV infections to precancers and cancers [2].

Detection of individual HPV genotypes associated with infections is important for research and, to some degree, for clinical use [3–5]. Epidemiologic research regarding the natural history of HPV depends on typing, because transmission, clearance, transformation, and invasive potential may differ by genotype. In a clinical setting, HPV testing is more efficient than the Pap test for primary cervical cancer screening [4, 6–9]. Evaluation of type-specific risk for precancer and cancer is necessary to decide which genotypes should be included in HPV assays and which genotypes should be detected individually. Recently approved assays use partial genotyping for HPV16 and HPV18 (and sometimes HPV45) for additional risk stratification [10]; more extended genotyping is under consideration.

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Correspondence: N. Wentzensen, MD, PhD, MS, Division of Cancer Epidemiology and Genetics, National Cancer Institute, 9609 Medical Center Drive, Room 6E448, Rockville, MD 20850 (wentzen@mail.nih.gov).

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Typing is important to estimate HPV vaccine efficacy (VE) [11]. Human papillomavirus vaccine trials and postvaccination surveillance programs require accurate and reliable HPV genotyping assays that detect a wide range of genotypes that are either (1) directly targeted by the vaccine, (2) targets of cross-protection, or (3) not directly or indirectly affected by vaccines [11]. There is a growing public health need for reliable, low-cost HPV genotyping methods applicable to large populations.

Few commercial assays are currently available for HPV genotyping, and most are laborious, low-throughput, and expensive. Next-generation sequencing (NGS) technology provides massively parallel sequencing capacity that can be used for detection of viral deoxyribonucleic acid (DNA) in clinical specimens. We recently developed TypeSeq, a novel NGS-based HPV genotyping assay that allows for highly automated, high-throughput testing at low cost [12]. In this study, we present the validation of TypeSeq in 2 large studies with a focus on detection of cervical precancers and on estimating VE.

## METHODS

### Study Descriptions

Human papillomavirus genotyping was performed in 2804 samples from women enrolled in the Study to Understand Cervical Cancer Early Endpoints and Determinants (SUCCEED) and in 2357 samples from women enrolled into the Costa Rica HPV Vaccine Trial (CVT). Of note, 1000 samples were randomly selected from each arm in CVT, and the remaining 357 samples were enriched for SPF-10 HPV16/18-positive samples.

This study is registered with Clinicaltrials.gov (NCT00128661). GlaxoSmithKline Biologicals provided vaccine and support for aspects of the trial associated with regulatory submission needs of the company under a Clinical Trials Agreement (FDA BB-IND 7920) during the 4-year, randomized, blinded phase of our study.

### SUCCEED Population, DNA Extraction, and Linear Array Genotyping

SUCCEED was a cross-sectional study of women 18 years of age or older with an abnormal Pap smear who were referred to colposcopy or treatment at the University of Oklahoma (OUHSC) between 2003 and 2011. Written informed consent was obtained from all women enrolled in the study, and Institutional Review Board approval was provided by OUHSC and the US National Cancer Institute (NCI) [13].

The DNA isolation method used for SUCCEED has been described previously [14]. Linear Array HPV Genotyping System (Roche Molecular Diagnostics) genotyping was performed as described previously [14–16]. Up to 80 patient specimens, 3 HPV16-positive controls, and 1 HPV-negative control were amplified in each batch using the Linear Array (LA). Detection of both  $\beta$ -globin concentration control probes was required to report genotyping results. A hybridization signal was called “positive” when an unambiguous, continuous band was observed on the array.

### CVT Population, DNA Extraction, and SPF10-LiPA Genotyping

The CVT was a community-based, double-blind, randomized, controlled phase III trial of the bivalent vaccine (Cervarix). As previously described, 18- to 25-year-old women residing in the provinces of Guanacaste and Puntarenas, Costa Rica, identified via a population census specifically conducted for the study, were invited to participate by attending a study clinic [17]. The trial was approved by human subjects review committees of the NCI and Instituto Costarricense de Investigación y Enseñanza en Nutrición y Salud ([INCIENSA] Costa Rica).

Extracted DNA from cervical specimens was polymerase chain reaction (PCR)-amplified and hybridized to HPV-specific probes using the SPF10 HPV DNA enzyme immunoassay (DEIA) system and the LiPA25, version 1, line detection system (SPF10 DEIA/LiPA25/TS16/18 system) [17, 18], following the manufacturer’s instructions. In addition, all specimens positive for HPV DNA using SPF10 DEIA but negative for HPV16 or HPV18 by LiPA25 were also tested with type-specific primers/probes for the presence of HPV16 and HPV18 DNA [18, 19].

### TypeSeq Genotyping

TypeSeq is able to detect the following 51 types: HPV3, 6, 11, 13, 16, 18, 26, 28, 30, 31, 32, 33, 34, 35, 39, 40, 42, 43, 44, 45, 51, 52, 53, 54, 56, 58, 59, 61, 62, 66, 67, 68, 69, 70, 71, 72, 73, 74, 76, 81, 82, 83, 84, 85, 86, 87, 89, 90, 91, 97, 114 (see [Supplementary Table 1](#)). Up to 950 patient specimens, 12 HPV-positive control pools, 2 HPV-negative human-positive controls, and 2 no-template controls were processed per batch, with 1 additional no-template control randomly located on each 96-well plate of specimens. Human beta-2-microglobulin (*B2M*) gene (GenBank accession number NG\_012920) served as the internal positive control in each reaction.

In brief, the stage 1 (S1) type-specific multiplex amplification and copy number standardization PCR was performed in a final reaction volume of 12  $\mu$ L, containing 5  $\mu$ L purified genomic DNA. The S1 primer pool contained 127 RNase H2-dependent primers (Integrated DNA Technologies, Coralville, IA), targeting 1 human gene (*B2M*) and a region of the L1 gene. The S2 primer pool contained nested *B2M* primers and 170 nested HPV unmodified primers (Integrated DNA Technologies). After cycling, unincorporated primers were degraded with Exonuclease I (Lucigen, Middleton, WI), then 2  $\mu$ L was used as template for the 10- $\mu$ L stage 3 (S3) PCR. During the S3 PCR, Ion sequencing adapters and dual barcodes were incorporated into amplicons via the universal priming sites. The PCR products were pooled and purified, then sequenced on the Ion S5 platform (Thermo Fisher Scientific, Waltham, MA) according to the manufacturer’s instructions. Dual barcode demultiplexing, quality filtering and HPV genotyping detection were performed using a custom plugin developed in-house, run within the Torrent Suite software (Thermo Fisher Scientific). A minimum of 850 total HPV reads or 300 *B2M* reads per sample was

required to report genotyping results, otherwise the sample was reported as “failed to amplify”. Positive HPV type calls required a minimum of 127–212 reads, depending on type.

Five hundred fifty-eight CVT specimens were tested in 2 independent batches on different days to assess TypeSeq interbatch reproducibility. Batch 1 was performed using manual pipetting, and batch 2 was done by automated pipetting using a JANUS liquid handler (PerkinElmer, Waltham, MA).

### Statistical Analysis

We calculated overall agreement and positive agreement between TypeSeq and LA genotyping in SUCCEED, and between TypeSeq and SPF10-LiPA genotyping in CVT, and we calculated McNemar *P* values. We further evaluated agreement between hierarchical categories of HPV positivity between the paired tests, combining any carcinogenic type (HR<sup>+</sup>), other HPV types, and types only detected by TypeSeq but not by the paired assay. We compared the observed frequencies of 2-type combinations for the 37 genotypes detected by LA and for the 51 genotypes detected by TypeSeq with expected frequencies in all women with at least 2 concurrent HPV infections. To obtain expected frequencies for a 2-type combination, the observed genotype frequencies for both types were multiplied and the result was multiplied with the total number of subjects.

In SUCCEED, we compared the risk of precancer and cancer (cervical intraepithelial neoplasia [CIN]3<sup>+</sup>) for each genotype as determined by TypeSeq and LA. For testing statistical significance, we compared the proportion of TS<sup>+</sup>/LA<sup>−</sup> and TS<sup>−</sup>/LA<sup>+</sup> samples that were positive for CIN3<sup>+</sup> using Fisher’s exact test. To evaluate the performance of the assay for HPV vaccination studies, we recomputed VE at the 4-year study visit in CVT using HPV results generated by TypeSeq and compared these estimates to those obtained in the initial round of testing using the SPF10 DEIA/LiPA25/TS16/18 system. Vaccine efficacy against one-time detection of HPV16/18, HPV31/33/45, and other carcinogenic HPV types (excluding HPV16, 18, 31, 33, and 45) based on the TypeSeq and SPF10 tests in the intention-to-treat cohort (all vaccinated women) was computed. For each arm, we defined the prevalence as the proportion of the number of events among the number of women at the 4-year study visit. The complement of the ratios of the HPV prevalence in the HPV arm and the control arm are the VE estimates. We calculated these VE estimates within the randomly sampled population and weighted back to the entire CVT. Exact confidence intervals (CIs) [20] for VE were calculated based on the binomial distribution of the number of events in the HPV arm among the total number of events in the HPV and control arms [21]. Vaccine efficacy obtained by the SPF10-LiPA system was predefined as the comparator. The prespecified goal of the analysis was that the 95% CI of the TypeSeq VE estimate included the SPF10-LiPA/DEIA VE point estimate, to assure comparability of the VE estimates between both tests.

## RESULTS

### Prevalence of 51 Human Papillomavirus Genotypes Detected by TypeSeq in SUCCEED and CVT

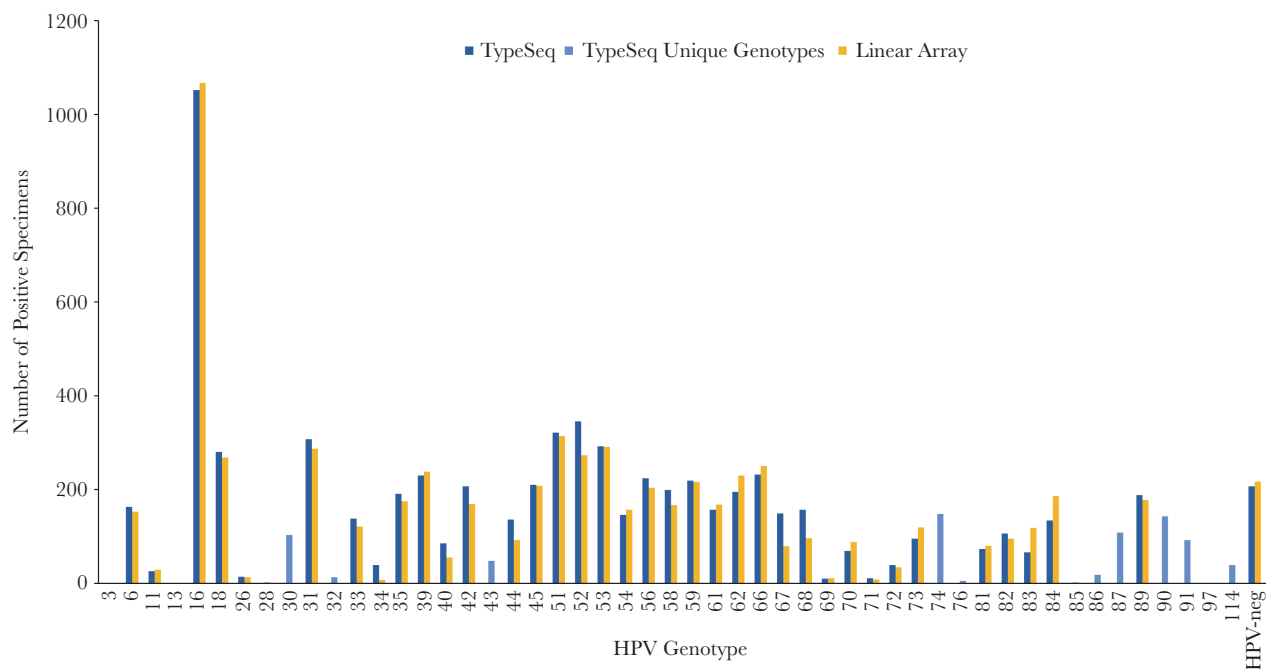
We evaluated TypeSeq results of cervical samples from 2804 women enrolled in SUCCEED and from 2357 women enrolled in CVT, respectively. In SUCCEED, a colposcopy referral population enriched for precancers and cancers, most women were HPV positive and HPV16 was by far the most common type (Figure 1). Among the 37 HPV genotypes detected by both TypeSeq and LA, the number of infections detected was very similar for both assays. The most common types detected by TypeSeq not included in LA were HPV30, 43, 74, 87, 90, 91, and 114; none has known strong disease associations. In CVT, an HPV vaccination trial of young women, a large proportion of women were negative for all 51 types detected using TypeSeq. For most of the 26 types detected by both TypeSeq and SPF10-LiPA, the number of infections was similar for both assays (Figure 2). To evaluate possible cross-reactivity between HPV genotypes, we compared observed versus expected numbers of 2-genotype combinations detected by LA and TypeSeq in SUCCEED. With LA, we observed substantially more combinations of HPV56 and HPV66, which are closely related and have been previously described to cross-react (see [Supplementary Figure 1A](#)) [15]. In contrast, there was no type combination observed substantially more frequently than expected for TypeSeq, suggesting no cross-reactivity (see [Supplementary Figure 1B](#)).

To assess TypeSeq reproducibility on clinical specimens, we tested 558 CVT specimens in duplicate using manual and automated processing (Table 1). Agreements for testing positive were 93.1% for any HR type, 93.2% for HPV16/18 combined, and ranged from 71.4% (HPV59) to 100% (HPV58) for individual HR types. Positive agreements were 60% or higher for 33 of the 35 LR types with positive specimens in either batch. None of the discrepancies were statistically significant for any type (*P* < .05, McNemar). Five types (HPV3, 13, 28, 76, and 97) were not detected in either batch.

### Agreement of TypeSeq With Linear Array and SPF10-LiPA for Individual Human Papillomavirus Types

We evaluated the agreement of TypeSeq with LA in SUCCEED and SPF10-LiPA in CVT. Comparing TypeSeq with LA in SUCCEED, 10 of 13 carcinogenic types had at least 80% positive agreement and HPV16 had 94% positive agreement (Table 2). The 3 types with lower agreement were HPV51, 52, and 56; all showed higher detection with TypeSeq. Six types (HPV31, 33, 35, 52, 56, and 58) showed a significantly higher positivity for TypeSeq among the discrepant results (McNemar, *P* < .05). There was high agreement for detecting overall HPV positivity, and positive agreement for detecting any HR type reached 95% between TypeSeq and LA (see [Supplementary Table 2](#)).

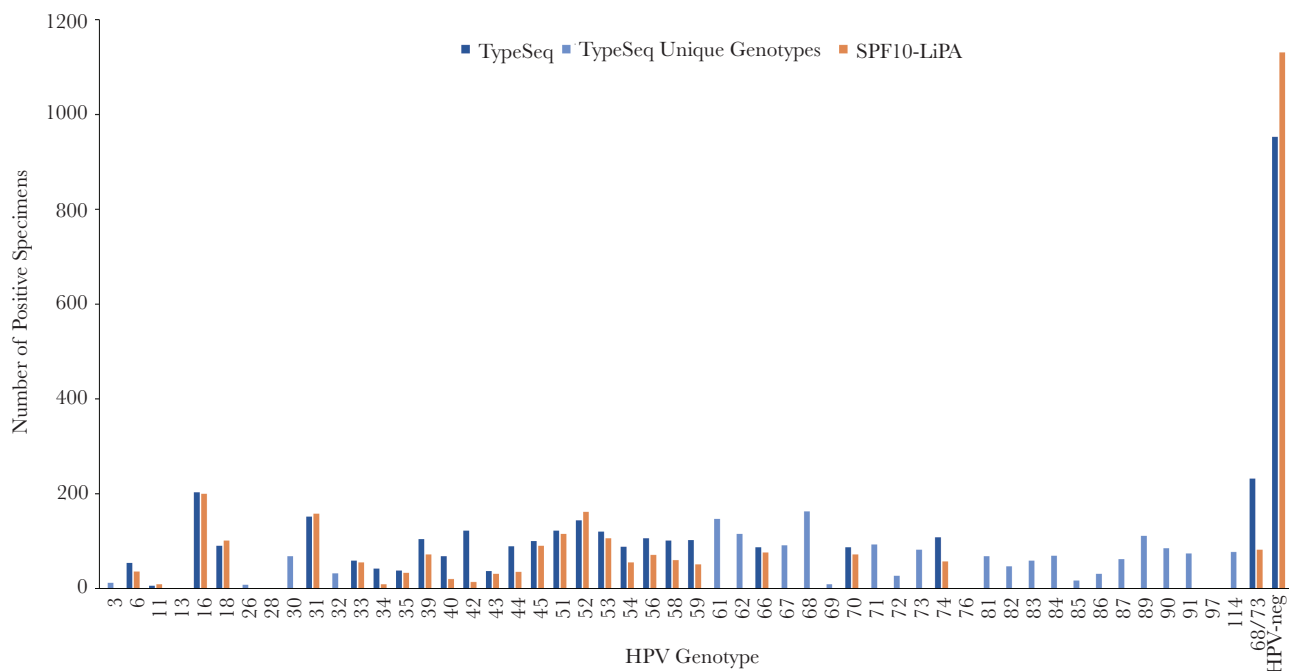
Positive agreement between TypeSeq and SPF10-LiPA was somewhat lower, with 4 types having at least 70% and 9 types



**Figure 1.** TypeSeq versus Linear Array (LA) positives by human papillomavirus (HPV) genotype for 2804 clinical specimens. The number of positive specimens are shown for LA and TypeSeq for the 37 genotypes detectable by both assays. The genotypes uniquely detectable by TypeSeq are displayed as “TypeSeq Unique Genotypes”. Linear Array’s HPV55 positives are shown as HPV44 according to the current PAVE classification. HPV68 represents results for the HPV68 lineages C to F (previously “68b”) detectable by LA. HPV82 represents a combined result for HPV82 and 82v (IS39), which are detected individually by both assays.

having at least 60% positive agreement (Table 3). The positive agreement for HPV16 reached 80%. The types with lower agreement were HPV35, 58, 59, and 68/73; all showed higher

detection by TypeSeq. Five types (HPV39, 56, 58, 59, and 68) showed a significantly higher positivity by TypeSeq among the discrepant results ( $P < .00001$ , McNemar), whereas HPV52



**Figure 2.** TypeSeq versus SPF10-LiPA positives by human papillomavirus (HPV) genotype for 2357 clinical specimens. The number of positive specimens are shown for SPF10-LiPA and TypeSeq for the 27 genotypes detectable by both assays. The genotypes uniquely detectable by TypeSeq are displayed as “TypeSeq Unique Genotypes”.

**Table 1. TypeSeq Reproducibility Testing on 558 CVT Clinical Specimens in Duplicate<sup>a</sup>**

HPV Genotype	Number of Specimens				%Agreement		
	R1 <sup>-</sup> /R2 <sup>-</sup>	R1 <sup>+</sup> /R2 <sup>-</sup>	R1 <sup>-</sup> /R2 <sup>+</sup>	R1 <sup>+</sup> /R2 <sup>+</sup>	Total	Positive	McNemar <i>P</i> Value
Any HPV	175	21	14	348	93.7	90.9	.31
Any HR-HPV <sup>b</sup>	313	12	5	228	97.0	93.1	.15
16/18	514	1	2	41	99.5	93.2	1
3	558	0	0	0	100.0	NE	NE
6	544	2	0	12	99.6	85.7	.48
11	557	0	0	1	100.0	100.0	NE
13	558	0	0	0	100.0	NE	NE
16	522	1	1	34	99.6	94.4	.48
18	548	0	1	9	99.8	90.0	1
26	555	0	0	3	100.0	100.0	NE
28	558	0	0	0	100.0	NE	NE
30	535	3	3	17	98.9	73.9	.68
31	531	1	2	24	99.5	88.9	1
32	548	2	0	8	99.6	80.0	.48
33	546	1	0	11	99.8	91.7	1
34	547	1	2	8	99.5	72.7	1
35	543	0	1	14	99.8	93.3	1
39	528	2	1	27	99.5	90.0	1
40	546	2	3	7	99.1	58.3	1
42	536	6	0	16	98.9	72.7	.041
43	550	3	2	3	99.1	37.5	1
44	533	3	5	17	98.6	68.0	.72
45	541	2	0	15	99.6	88.2	.48
51	533	2	5	18	98.7	72.0	.45
52	510	2	1	45	99.5	93.8	1
53	529	4	3	22	98.7	75.9	1
54	541	2	4	11	98.9	64.7	.68
56	532	1	0	25	99.8	96.2	1
58	525	0	0	33	100.0	100.0	NE
59	530	7	1	20	98.6	71.4	.077
61	534	0	3	21	99.5	87.5	.25
62	534	1	3	20	99.3	83.3	.62
66	528	1	0	29	99.8	96.7	1
67	536	0	4	18	99.3	81.8	.13
68a	527	4	4	23	98.6	74.2	.72
68b	549	1	0	8	99.8	88.9	1
69	556	0	0	2	100.0	100.0	NE
70	533	2	3	20	99.1	80.0	1
71	540	0	1	17	99.8	94.4	1
72	551	0	2	5	99.6	71.4	.48
73	546	2	2	8	99.3	66.7	.62
74	528	6	2	22	98.6	73.3	.28
76	558	0	0	0	100.0	NE	NE
81	540	3	0	15	99.5	83.3	.25
82	555	1	0	2	99.8	66.7	1
82v	547	1	1	9	99.6	81.8	.48
83	545	2	1	10	99.5	76.9	1
84	543	0	1	14	99.8	93.3	1
85	552	0	1	5	99.8	83.3	1
86	549	1	1	7	99.6	77.8	.48
87	535	3	2	18	99.1	78.3	1
89	538	4	1	15	99.1	75.0	.37
90	543	1	3	11	99.3	73.3	.62
91	543	0	0	15	100.0	100.0	NE
97	558	0	0	0	100.0	NE	NE
114	543	3	3	9	98.9	60.0	.68

Abbreviations: CVT, Costa Rica Vaccine Trial; HPV, human papillomavirus; HR-HPV, high-risk HPV; NE, not evaluable; R1, replicate 1; R2, replicate 2.

<sup>a</sup>Clinical specimens were tested in duplicate by manual (R1) or automated (R2) processing.

<sup>b</sup>HR-HPV represents HPV types 16, 18, 31, 33, 35, 39, 45, 51, 52, 56, 58, 59, and 68.



**Table 2. Agreement Between TypeSeq and Linear Array for 13 Carcinogenic Types Among 2804 Women in SUCCEED**

HPV Type	N LA	% LA	N TypeSeq	% TypeSeq	N Both <sup>+</sup>	% Both <sup>+</sup>	N LA <sup>+</sup> /TypeSeq <sup>-</sup>	% LA <sup>+</sup> /TypeSeq <sup>-</sup>	N LA <sup>-</sup> /TypeSeq <sup>+</sup>	% LA <sup>-</sup> /TypeSeq <sup>+</sup>	N Neither <sup>+</sup>	% Neither <sup>+</sup>	% Agreement	% Positive Agreement	McNemar PValue
16	1067	38.1	1052	37.6	1025	36.6	42	1.5	27	1.0	1707	60.9	97.5	93.7	.09
18	268	9.6	280	10.0	250	8.9	18	0.6	30	1.1	2503	89.4	98.3	83.9	.11
31	287	10.3	307	11.0	275	9.8	12	0.4	32	1.1	2482	88.6	98.4	86.2	.003
33	121	4.3	138	4.9	120	4.3	1	0.04	18	0.6	2662	95.0	99.3	86.3	.00004
35	175	6.3	191	6.8	167	6.0	8	0.3	24	0.9	2602	92.9	98.9	83.9	.004
39	238	8.5	230	8.2	216	7.71	22	0.8	14	0.5	2549	91.0	98.7	85.7	.24
45	208	7.4	210	7.5	189	6.75	19	0.7	21	0.8	2572	91.8	98.6	82.5	.87
51	314	11.2	321	11.5	266	9.5	48	1.7	55	2.0	2432	86.8	96.3	72.1	.55
52	273	9.7	345	12.3	250	8.92	23	0.8	95	3.4	2436	86.9	95.8	67.9	.0002
56	204	7.3	224	8.0	184	6.57	20	0.7	40	1.4	2557	91.3	97.9	75.4	.01
58	167	6.0	199	7.1	163	5.82	4	0.1	36	1.3	2598	92.8	98.6	80.3	<.00001
59	216	7.7	219	7.8	197	7.03	19	0.7	22	0.8	2563	91.5	98.5	82.8	.76
68 <sup>a</sup>	96	3.4	103	3.7	90	3.21	6	0.2	13	0.5	2692	96.1	99.3	82.6	.17

Abbreviations: HPV, human papillomavirus; LA, Linear Array; SUCCEED, Study to Understand Cervical Cancer Endpoints and Early Determinants.

<sup>a</sup>HPV68 represents results for the lineages detectable by LA (C to F, formerly "68b"). TypeSeq results for HPV68 lineages A and B (formerly "68a"), which are uniquely detectable by TypeSeq, were excluded from this analysis.

was significantly more positive in SPF10-LiPA ( $P = .03$ ). The positive agreement between TypeSeq and SPF10-LiPA in the random draw of 1998 samples from both study arms was slightly lower for the types that were oversampled (see [Supplementary Table 3](#)). Positive agreement for detecting any HR type reached 77% between TypeSeq and SPF10-LiPA (see [Supplementary Table 4](#)).

#### Human Papillomavirus Genotype Prevalence by TypeSeq and Linear Array in CIN3<sup>+</sup> From SUCCEED

We compared the genotype prevalence in cervical precancer and cancer (CIN3<sup>+</sup>) by TypeSeq and LA ([Table 4](#)). Overall, there was high concordance between both assays for all genotypes. Cases with HPV39, 52, 56, and 58 showed a difference of at least 5 CIN3<sup>+</sup> cases between both assays; TypeSeq detected more CIN3<sup>+</sup> for all these types except for

HPV39-positive cases, in which it detected fewer cases of CIN3<sup>+</sup>.

When comparing all TypeSeq-positive and all LA-positive women, the risk of CIN3<sup>+</sup> was very similar for all genotypes. There was no significant difference in the risk of CIN3<sup>+</sup> between the discrepant categories (TS<sup>-</sup>/LA<sup>+</sup> vs TS<sup>+</sup>/LA<sup>-</sup>) for any genotype ( $P > .05$ ).

#### Estimation of Vaccine Efficacy Based on TypeSeq Compared With SPF10-LiPA Test Results

To evaluate the performance of TypeSeq for detecting viral endpoints in vaccine trials and population surveillance studies, we calculated vaccine efficiency of the bivalent HPV vaccine based on HPV genotyping with TypeSeq compared with the reference standard in CVT, SPF10-LiPA ([Table 5](#)). Vaccine efficiency estimates were computed in a population naive to HPV

**Table 3. Agreement Between TypeSeq and SPF10-LiPA for 13 Carcinogenic Types Among 2357 Women in CVT**

HPV Type	N SPF10	% SPF10	N TypeSeq	% TypeSeq	N Both <sup>+</sup>	% Both <sup>+</sup>	N SPF10 <sup>+</sup> /TypeSeq <sup>-</sup>	% SPF10 <sup>+</sup> /TypeSeq <sup>-</sup>	N SPF10 <sup>-</sup> /TypeSeq <sup>+</sup>	% SPF10 <sup>-</sup> /TypeSeq <sup>+</sup>	N Neither <sup>+</sup>	% Neither <sup>+</sup>	% Agreement	% Positive Agreement	McNemar PValue
16	200	8.5	203	8.6	179	7.6	21	0.89	24	1.0	2133	90.5	98.1	79.9	.38
18	101	4.3	90	3.8	84	3.6	17	0.72	6	0.3	2250	95.5	99.0	78.5	.43
31	159	6.8	152	6.5	125	5.3	34	1.44	27	1.2	2171	92.1	97.4	67.2	.44
33	55	2.3	59	2.5	46	2.0	9	0.38	13	0.6	2289	97.1	99.1	67.7	.39
35	33	1.4	38	1.6	25	1.1	8	0.34	13	0.6	2311	98.1	99.1	54.4	.28
39	72	3.1	104	4.4	66	2.8	6	0.25	38	1.6	2247	95.3	98.1	60.0	<.00001
45	90	3.8	100	4.2	80	3.4	10	0.42	20	0.9	2247	95.3	98.7	72.7	.07
51	115	4.9	122	5.2	96	4.1	19	0.81	26	1.1	2216	94.0	98.1	68.1	.30
52	163	6.9	144	6.1	120	5.1	43	1.82	24	1.0	2170	92.1	97.2	64.2	.03
56	71	3.0	106	4.5	69	2.9	2	0.08	37	1.6	2249	95.4	98.4	63.9	<.00001
58	60	2.6	101	4.3	58	2.5	2	0.08	43	1.8	2254	95.6	98.1	56.3	<.00001
59	51	2.2	102	4.3	49	2.1	2	0.08	53	2.3	2253	95.6	97.7	47.1	<.00001
68/73	83	3.5	232	9.8	78	3.3	5	0.21	154	6.5	2120	89.9	93.3	32.9	<.00001

Abbreviations: CVT, Costa Rica Vaccine Trial; HPV, human papillomavirus.

**Table 4. Prevalence of HPV Genotypes in CIN3<sup>+</sup> by TypeSeq and Linear Array**

HPV Type	TS <sup>+</sup>	TS <sup>+</sup> CIN3 <sup>+</sup>	Risk TS (%)	LA <sup>+</sup>	LA <sup>+</sup> CIN3 <sup>+</sup>	Risk LA (%)	TS <sup>-</sup> /LA <sup>-</sup>	TS <sup>-</sup> /LA <sup>+</sup> CIN3 <sup>+</sup>	Risk TS <sup>-</sup> /LA <sup>+</sup> a (%)	TS <sup>+</sup> /LA <sup>-</sup>	TS <sup>+</sup> /LA <sup>-</sup> CIN3 <sup>+</sup>	Risk TS <sup>+</sup> /LA <sup>-</sup> a (%)	TS <sup>+</sup> /LA <sup>+</sup> CIN3 <sup>+</sup>	Risk TS <sup>+</sup> /LA <sup>+</sup> (%)
16	1052	474	45.1	1067	476	44.6	1707	42	5	11.9	27	3	11.1	46.0
18	280	75	26.8	268	74	27.6	2503	18	6	33.3	30	7	23.3	27.2
31	307	81	26.4	287	78	27.2	2482	12	2	16.7	32	5	15.6	27.6
33	138	39	28.3	121	35	28.9	2662	1	0	0.0	18	4	22.2	29.2
35	191	37	19.4	175	33	18.9	2602	8	2	25.0	24	6	25.0	18.6
39	230	27	11.7	238	33	13.9	2549	22	7	31.8	14	1	7.1	12.0
45	210	56	26.7	208	60	28.8	2572	19	9	47.4	21	5	23.8	27.0
51	321	43	13.4	314	40	12.7	2432	48	8	16.7	55	11	20.0	12.0
52	345	63	18.3	273	54	19.8	2436	23	7	30.3	95	16	16.8	18.8
56	224	28	12.5	204	21	10.3	2557	20	2	10.0	40	9	22.5	10.3
58	199	42	21.1	167	34	20.4	2598	4	0	0.0	36	8	22.2	20.9
59	219	38	17.4	216	39	18.1	2563	19	4	21.1	22	3	13.6	17.8
68	103	21	20.4	96	19	19.8	2692	6	0	0.0	13	2	15.4	21.1

Abbreviations: CIN, cervical intraepithelial neoplasia; HPV, human papillomavirus; LA, Linear Array; TS, TypeSeq.

\*The difference in risk between TS<sup>-</sup>/LA<sup>-</sup> and TS<sup>+</sup>/LA<sup>+</sup> was not significant for any of the types ( $P > .5$ ).

infections at the time of vaccination as well as the full analytic cohort for HPV16/18 (the types included in the bivalent vaccine), HPV31, 33, 45 (3 types that show cross-protection for the bivalent vaccine), and 7 other types that are typically not affected by the vaccine. The number of infections detected by both assays was similar. In all comparisons, there was no statistically significant difference in VE as determined by TypeSeq compared with SPF10-LiPA, and the prespecified threshold for assay comparability was achieved in all population and HPV genotype groups.

## DISCUSSION

TypeSeq is a novel, affordable, high-throughput NGS assay for detection of 51 HPV genotypes. We validated TypeSeq in over 5100 cervical specimens from 2 large epidemiological studies. TypeSeq showed high agreement with 2 widely established HPV genotyping assays, LA and SPF10-LiPA. In a large study of cervical precancers, no difference in risk of CIN3<sup>+</sup> associated with individual genotypes was observed between TypeSeq and LA. In a randomized controlled HPV vaccine trial, no difference in VE for vaccine types and cross-protective types was observed between TypeSeq and SPF10-LiPA.

Human papillomavirus genotyping is central to understanding natural histories of HPV-related diseases and to HPV-based prevention efforts [1, 22]. Natural history differs between genotypes with respect to HPV acquisition and progression to precancer [23, 24]. Most importantly, the risk of cancer differs substantially between individual genotypes [2]. However, currently, very few HPV genotyping assays are commercially available, and most established assays are costly and laborious, which has limited HPV genotyping particularly in large longitudinal population-based studies with multiple rounds of specimen sampling. More important, the performance of HPV assays needs to strike a delicate balance

between sensitivity and specificity, so that all clinically important infections, but not irrelevant minor infections or viral depositions, are detected.

We are now entering a new era of HPV-based prevention of cervical cancers, with 3 highly efficacious HPV vaccines available that cover different HPV types. Evaluation of VE is shifting to viral endpoints, which requires accurate and reproducible HPV genotyping [11]. A new focus of HPV vaccine studies evaluates efficacy and effectiveness of vaccination with fewer doses, particularly 1-dose vaccination [25]. Likewise, postvaccination surveillance requires affordable and reliable HPV genotyping of specimens from large populations.

At the same time, HPV genotyping is important to determine which types to include in HPV assays and which types to detect individually for additional risk stratification [1]. Given its comparable performance to LA, an assay that performs similarly to HPV assays approved for HPV screening [26], TypeSeq could also be evaluated as an assay for screening and triage.

TypeSeq can process up to 950 samples per batch. The turnaround time for this batch size is 3 days, with a hands-on time of 12 hours. Human papillomavirus genotype calling is fully automated, and results are immediately exported into a spreadsheet. All genotypes are reported individually, but the software can mask and group genotypes, allowing customization of the assay to specific needs.

To improve detection of HPV genotypes in multiple infections, the assay normalizes the input viral DNA over a wide range of concentrations. Therefore, the assay does not provide information about viral load. However, this is a minor limitation since the clinical relevance of viral load is limited [27].

The unique features of TypeSeq, particularly the high throughput, low cost, and high accuracy, make it an attractive assay for many applications. It can provide reliable HPV genotyping for large natural history studies with multiple



**Table 5. Bivalent Vaccine Efficacy Determined by TypeSeq and SPF10-LiPA**

HPV16/18				HPV31, 33, 45				Other (HPV35, 39, 51, 52, 56, 58, 59)					
Assay	Trial Arm	ATP		ITT		ATP		ITT		ATP		ITT	
		Events/ Women	VE (%) (95% CI)	Events/ Women	VE (%) (95% CI)	Events/ Women	VE (%) (95% CI)	Events/ Women	VE (%) (95% CI)	Events/ Women	VE (%) (95% CI)	Events/ Women	VE (%) (95% CI)
All Women Tested by TypeSeq													
SPF10-LiPA	HPV	49/820	66.6	62/1104	69.2	69/823	45.7	99/1104	39.4	153/824	12.7	208/1104	9.8
	HAV	164/916	(54.3–75.9)	228/1250	(59.4–76.9)	142/920	(27.8–59.4)	185/1250	(22.8–52.7)	196/922	(–7.9 to 29.4)	261/1250	(–8.2 to 24.8)
TypeSeq	HPV	43/820	70.2	60/1104	69.5	66/823	52.7	90/1104	49.6	182/824	13.7	246/1104	7.8
	HAV	161/916	(58.5–78.9)	223/1250	(59.7–77.2)	156/920	(37.1–64.7)	202/1250	(35.5–60.8)	236/922	(–4.7 to 28.9)	302/1250	(–9.1 to 22.1)
Random Sample From Both Arms													
SPF10-LiPA	HPV	16/741	72.6	20/998	74.0	20/742	64.4	32/998	53.5	127/743	1.2	175/998	–6.3
	HAV	58/735	(53.2–84.7)	77/1000	(58.0–84.4)	56/739	(41.4–79.1)	69/1000	(29.7–69.8)	128/740	(–26.4 to 22.7)	165/1000	(–31.5 to 14.1)
TypeSeq	HPV	17/741	73.2	26/998	69.0	23/742	67.3	35/998	58.2	153/743	4.2	212/998	–8.4
	HAV	63/735	(55.0–84.7)	84/1000	(52.3–80.3)	70/739	(48.1–79.9)	84/1000	(38.4–72.1)	159/740	(–19.7 to 23.3)	196/1000	(–31.7 to 10.8)
Random Sample Weighted Back to Full Study													
SPF10-LiPA	HPV	49/2379	70.4	62/3175	72.7	69/2384	51.8	100/3175	46.3	385/2387	2.7	516/3175	0.3
	HAV	164/2340	(61.6–77.2)	228/3167	(66.3–77.9)	142/2352	(39.7–61.4)	185/3167	(37.0–54.2)	390/2354	(–19.9 to 21.1)	516/3167	(–19.5 to 16.8)
TypeSeq	HPV	55/2379	67.9	82/3175	65.7	82/2384	55.9	112/3175	52.6	460/2387	6.0	623/3175	–2.8
	HAV	169/2340	(54.6–77.4)	239/3167	(54.0–74.5)	184/2352	(40.9–67.1)	236/3167	(39.5–62.8)	483/2354	(–13.2 to 22.1)	605/3167	(–21.0 to 12.7)

Abbreviations: ATP, according-to-protocol; CI, confidence interval; HAV, control arm; HPV, vaccine arm; ITT, intention-to-treat; VE, vaccine efficacy.

sampling. We successfully evaluated TypeSeq in a randomized controlled vaccine trial, and, based on its performance, we plan to use it in future HPV vaccination studies. The performance of TypeSeq is similar to LA, a widely evaluated assay that has high agreement with clinical HPV tests like Cobas and Onclarity [28]. Further evaluation of TypeSeq as a clinical test for primary screening with partial genotyping is supported by our findings. Evaluation of other specimen types, such as anal and oral swab samples, as well as tissue specimens, using the assay is currently underway.

TypeSeq requires an infrastructure suitable for NGS technology and investment in sequencing equipment. However, these technologies are advancing quickly, and new robust equipment can now be made available in all regions of the world. Furthermore, smaller and more achievable NGS platforms are now available that reduce up-front costs. Due to the highly integrated and automated workflow of the assay, technology transfer is feasible and is currently happening in other laboratories.

## CONCLUSIONS

In summary, we have validated a novel NGS-based HPV genotyping assay that addresses a widespread need for high-throughput, affordable HPV genotyping for research, surveillance, and clinical management.

## Supplementary Data

Supplementary materials are available at *The Journal of Infectious Diseases* online. Consisting of data provided by the authors to benefit the reader, the posted materials are not copyedited and are the sole responsibility of the authors, so questions or comments should be addressed to the corresponding author.

## Notes

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**Potential conflicts of interest.** John T. Schiller and Douglas R. Lowy report that they are named inventors on US Government-owned HPV vaccine patents that are licensed to GlaxoSmithKline and Merck and for which the National Cancer Institute receives licensing fees. They are entitled to limited royalties as specified by federal law. All authors have submitted the ICMJE Form for Disclosure of Potential Conflicts of Interest. Conflicts that the editors consider relevant to the content of the manuscript have been disclosed.

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